- 1 Satellite Observations of Mexico City Pollution Outflow from the Tropospheric Emissions
- 2 Spectrometer (TES)
- 3 Changsub Shim¹, Qinbin Li^{1,2}, Ming Luo¹, Susan Kulawik¹, Helen Worden³, John Worden¹,
- 4 Annmarie Eldering¹, Glenn Diskin⁴, Glen Sachse⁴, Andy Weinheimer³, David Knapp³, Deedee
- 5 Montzca³, and Teresa Campos³
- 6
- ¹Jet Propulsion Laboratory, California Institute of Technology, CA
- 8 ²Department of Atmospheric & Oceanic Sciences, University of California, Los Angeles, CA
- 9 ³National Center for Atmospheric Research (NCAR), CO
- 10 ⁴NASA Langley Research Center, VA
- 11
- 12 Correspondence author:
- 13 Changsub Shim; cshim@jpl.nasa.gov
- 14

15

- Abstract
- 16 Concurrent tropospheric O₃ and CO vertical profiles from the Tropospheric Emission
- 17 Spectrometer (TES) during the MILAGRO/INTEX-B aircraft campaigns over the Mexico City
- 18 Metropolitan Area (MCMA) and its surrounding regions were used to examine Mexico City
- 19 pollution outflow on a regional scale. The pollution outflow from the MCMA occurred
- predominantly at 600-800 hPa as evident in O₃, CO, and NO_x enhancements in the *in situ* aircraft
- observations. TES O₃ and CO are sensitive to the MCMA pollution outflow due to their
- relatively high sensitivities at 600-800 hPa. We examined O₃, CO, and their correlation at 600-
- 23 800 hPa from TES retrievals, aircraft measurements, and GEOS-Chem model results. TES

captures much of the spatial and day-to-day variability of O_3 seen in the *in situ* data. TES CO, however, shows much less spatial and day-to-day variability compared with the *in situ* observations. The $\Delta O_3/\Delta CO$ slope is significantly higher in the TES data (0.43) than the *in situ* data (0.28) due partly to the lack of variability in TES CO. Extraordinarily high $\Delta O_3/\Delta CO$ slope (0.81) from TES observations at 618 hPa over the Eastern U.S. was previously reported by *Zhang et al.* (2006). Thus the application of TES CO-O₃ correlation to map continental pollution outflow needs further examination.

32 Keyword: Mega-city pollution, TES, O₃-CO correlation, INTEX-B, Mexico city

1. Introduction

Mega-city pollution has been a major environmental concern in developing and developed countries alike (*Fuchs et al.*, 1994). Pollution from these mega-cities has grave implications on regional to global air quality and climate change (*Mage et al.*, 1996; *Molina and Molina*, 2004). One of the world's most densely populated mega-cities, the Mexico City Metropolitan Area (MCMA: ~19°N, ~99°W, ~750 hPa), with a population of over 19 million within an area of ~1500 km², is among the most polluted cities (*IPCC*, 2007). Large emissions of air pollutants from the MCMA result from pervasive incomplete combustion including low-temperature household cooking and heating and from vehicle emissions (*CAM*, 2001). In 1998, for example, the total volatile organic compounds (VOCs) emissions from the MCMA (475 Gg/yr) is higher than that from Los Angeles (362 Gg/yr in 2001), the most polluted mega-city in the United Sates (*Molina and Molina*, 2002). The MCMA is located in the Valley of Mexico, a large basin in the high plateaus at the center of Mexico, at an altitude of 2.2 km above sea level, surrounded by

mountains on all four sides except a small opening at the north. The basin is often under the influence of high pressure systems, whose weak winds and persistent boundary layer thermal inversions often trap pollution to within the basin (*Molina and Molina*, 2002).

The MILAGRO/INTEX-B aircraft campaigns in March 2006 aimed to investigate the emissions, transport, and physiochemical transformation of key air pollutants from the MCMA (www.joss.ucar.edu/milagro/; Singh et al., 2008). During this campaign, the synoptic scale export of the MCMA pollution out of the Valley of Mexico is generally controlled by thermal-driven circulations in the boundary layer and southwesterly wind in the middle troposphere (Fast et al., 2007). Airborne measurements of various gaseous and aerosol pollutants were made aboard the NASA DC-8 and NSF C-130 aircrafts. In coordination with the aircraft observations, the Tropospheric Emission Spectrometer (TES) instrument aboard the Aura satellite conducted several special observations and global surveys (see Section 2.2) during the campaigns. TES provides concurrent global three-dimensional (3-D) mapping of tropospheric O₃ and CO (Beer, 2006).

Some recent studies have explored the use of space-borne observations to examine regional to global air pollution (*Boersma et al.*, 2008; *Clerbaux et al.*, 2008; *Zhang et al.*, 2006; *Massie et al.*, 2006). For example, *Zhang et al.* (2006) has shown that TES tropospheric data can be used to map global pollution and continental outflow. They found positive O_3 -CO correlations and slopes ($\Delta O_3/\Delta CO$) at 618 hPa over continental outflow regions, indicating significant photochemical O_3 production from pollution. In this study, we intend to examine TES observations of pollution outflow from the MCMA during MILAGRO/INTEX-B. The *in situ* aircraft observations allow us to evaluate TES O_3 and CO retrievals, in particular the spatial and day-to-day variability, and O_3 -CO correlations. CO is a good tracer for industrial and biomass

burning pollution (*Logan et al.*, 1981). The O₃-CO relationship, among other tracer-tracer correlations, has been used for ozone source attribution. It provides a reliable characterization of continental pollution outflow (*Fishman and Seiler*, 1983; *Chameides et al.*, 1987; *Parrish et al.*, 1993). Additionally, the O₃-CO correlation has also been used to evaluate the photochemical O₃ production in chemical transport models (*Chin et al.*, 1994).

We examine TES O_3 and CO and their correlations over the MCMA and surrounding regions against the aforementioned *in situ* aircraft observations. In particular, we focus on the spatial and temporal variability and $\Delta O_3/\Delta CO$ enhancement ratio. Results from the GEOS-Chem global 3-D chemical transport model are also compared with both the TES data and aircraft observations. We describe the aircraft measurements, TES retrievals, and GEOS-Chem in section 2. Time series of O_3 and CO and O_3 -CO correlations are discussed in section 3. Conclusions are given in section 4.

2. Methodology

84 2.1. Aircraft measurements

A suite of chemical tracers including O_3 , CO, NO_x , and VOCs were measured aboard the C-130 and DC-8 during the MILAGRO/INTEX-B campaigns. Figure 1 (left panel) shows the aircraft flight tracks. The C-130 flights stayed closer to the MCMA compared with those of the DC-8 with the later covering a much larger domain. O_3 and CO were measured by ChemiLuminescence Detector (CLD) and Tunable Diode Laser (TDL), respectively (*Avery et al.*, 2001; *Madronich et al.*, 2004; *Sachse et al.*, 1987) on the both C-130 and DC-8. The precisions of O_3 and CO are 3-7% and 1%, respectively with $\pm 5\%$ accuracy (*Richard et al.*, 2007; *Sachse*

et al., 1987; Singh et al., 2008). We used the 1-minute merged data. Our analysis focuses on observations from March 4-31, 2006 within 15-25°N and 95-103°W (rectangle, Figure 1).

2.2. TES Data

92

93

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

TES measures infrared emissions with high spectral resolution (0.1 cm⁻¹) and a wide spectral range from $660 - 2260 \text{ cm}^{-1}$ (Beer et al., 2001). TES overpass times (ascending node) are 01:45 and 13:45 local time. TES has a footprint of ~5×8 km in nadir, about 180 km apart between consecutive measurements along the orbital tracks in the standard Global Survey mode. Global coverage is achieved in 16 days. In the Step and Stare special observation mode, TES provides denser nadir spatial coverage (~40 km apart between consecutive measurements along the orbit) over a 60° latitudinal range (Beer et al., 2006). We use here O₃ and CO data from all eight Step and Stares and four global surveys from March 2006. The corresponding TES orbital tracks over the MCMA region during MILAGRO/INTEX-B are shown in Figure 1 (right panel). TES retrievals of O₃ and CO are based on the optimization between modeled a priori and observed radiance with maximum a posteriori method (Rodgers, 2000) and the details of the retrieval algorithms are described in Worden et al. (2004), Bowman et al. (2002, 2006), and Luo et al. (2007a). Here we use the version 3 data (V003, F04 04) (Osterman et al., 2006) available at the time of this study. The degrees of freedom for signal (DOFs) for O₃ and CO are 1.6 and 1.4, respectively in the troposphere (Worden et al., 2004; Bowman et al., 2006; Luo et al., 2007b). TES sensitivity is reduced by $\sim 40\%$ with a cloud optical depth of 0.1 in the lower troposphere (Kulawik et al., 2006). We excluded TES retrievals when cloud optical depth is larger than 0.2. We also excluded TES data with low DOFs (< 0.4). The validation of TES data with in situ measurements requires applying TES averaging kernels to in situ profiles to take into account the relatively coarse TES vertical resolution and a

priori information (Rodgers and Connor, 2003; Worden et al., 2007). On average, TES O₃ shows a positive bias of ~6 ppbv (15%) throughout the troposphere in comparison with ozonesonde data (Nassar et al., 2008; Worden et al., 2007). Richard et al. (2008) compared TES O₃ data with airborne in situ measurements during INTEX-B and found a positive bias of ~2-6 ppbv (5 – 15%) in the troposphere. TES CO has been validated with aircraft data during the Aura Validation Experiments (AVEs) by Lopez et al. (2008). They found that TES CO is slightly lower than the *in situ* data (< 10% or ~8 ppbv) in the middle latitudes and slightly higher (5 – 10% or ~4-8 ppbv) in the tropics. Luo et al. (2007) conducted TES CO validations during INTEX-B and showed consistent results (a negative bias of < 10%). Due to the lack of concurrent TES and in situ measurements, relaxed coincident criteria such as within ~500 km distance and ~48 hours were used in the aforementioned validations. Figure 2 shows the closest TES orbital track (a Global Survey) to MCMA on March 19th, 2006 (left panel) and averaging kernels for O₃ and CO (upper and lower right panels) along the orbital track between 15°N and 25°N over Mexico. The closest distance between the MCMA and this TES orbital track is 95 km at location "a" (Figure 2) over the Mexico City Basin (~20°N, \sim 750 hPa) where there is relatively poor sensitivity for both O₃ and CO. The reason for the weak sensitivity is under further investigation by the TES retrieval team, (K. Bowman, personal communication, 2008, Jet Propulsion Laboratory). By contrast, O₃ and CO averaging kernels away from the Mexico City Basin (~23.3°N, ~850 hPa; location "b" in Figure 2) show largest sensitivities to 600 – 800 hPa, typical for the middle and low troposphere under clear sky (e.g., Jones et al., 2003; Luo et al., 2007; Richard et al., 2008; Worden et al., 2007; Zhang et al., 2006). To illustrate the latitudinal variations of TES O₃ and CO averaging kernels, the diagonals of the TES O₃ and CO averaging kernels from the March 12th TES Step & Stares are shown in

115

116

117

118

119

120

121

122

123

124

125

126

127

128

129

130

131

132

133

134

135

136

137

Figure 3 for $5^{\circ}\text{S}-35^{\circ}\text{N}$. The closest distance from the orbital track to the MCMA is ~150 km. While there is relatively large sensitivity in the lower to middle troposphere, there is poor sensitivity for both O_3 and CO over the Mexico City Basin (17°–20°N). Thus it appears that TES data are sensitive to the MCMA pollution outflow, rather than the pollution right over Mexico City Basin itself.

2.3. GEOS-Chem

GEOS-Chem is a global 3-D chemical transport model driven by assimilated meteorological data from NASA Global Modeling Assimilation Office (GMAO) (*Bey et al.*, 2001). We use version 7-04-10 (http://www.as.harvard.edu/chemistry/trop/geos) of the model, driven by GEOS-4 meteorological data with a horizontal resolution of 2°×2.5° and 30 vertical layers. The meteorological fields are updated every six hours, and the surface fields and mixing depths are updated every three hours.

GEOS-Chem includes a comprehensive tropospheric O₃-NO_x-VOC chemistry mechanism. Climatological monthly mean biomass burning emissions are from *Duncan et al.* (2003). The fossil fuel emissions are from the Emission Database for Global Atmospheric Research (EDGAR) inventory for NO_x, CO, and SO₂ and from the Global Emission Inventory Activity (GEIA) for other chemical compounds (*Benkovitz et al.*, 1996; *Olivier et al.*, 2001). These emissions are updated with particular national emission inventories and fuel use data: the Big Bend Regional Aerosol and Visibility Observational Study (BRAVO) inventory for Mexico (*Kuhns et al.*, 2005) and U.S. EPA NEI 99 inventory (National Emissions Inventory, base year 1999, version 3) for the continental U.S. (*EPA*, 2004). The biogenic VOCs emissions are based on the Model of Emissions of Gases and Aerosols from Nature (MEGAN) inventory (*Guenther et al.*, 2006). The lightning NO_x emissions are parameterized based on cloud top height and

regionally scaled to climatological satellite observations of flash rates (*Hudman et al.*, 2007). We conducted GEOS-Chem simulations for September 2005-March 2006 with the first six months for initialization. We focus our analysis on March 2006. Model results are sampled along the aircraft flight tracks and TES orbital tracks.

To compare GEOS-Chem results with TES retrievals, model profiles of O₃ and CO are convoluted with TES averaging kernels to account for the different sensitivities and *a priori* information of TES retrievals to different pressure levels (*Jones et al.*, 2003; *Richards et al.*, 2008). The resulting model profiles can then be directly compared with TES retrievals (*Zhang et al.*, 2006; *Jourdain et al.*, 2007; *Worden et al.*, 2007).

We note here that the emissions used in our GEOS-Chem simulations are not 2006-specific. Additionally, the horizontal resolution of 2°×2.5° may be too coarse for resolving details of the regional pollution outflow. GEOS-Chem simulations with 1°×1° nested simulations over North America have been done before (e.g., *Li et al.*, 2005) but not available for 2006 at the time of this study. We include GEOS-Chem results for comparison despite the aforementioned shortcomings in our model simulations. Improving model simulations for the MILAGRO/INTEX-B period is obviously required to understand the processes controlling the MCMA outflow. It is however beyond the scope of the present analysis.

180 3. Results and discussions

3.1. Mexico City pollution outflow

The median vertical profiles of O_3 , CO, and NO_x mixing ratios from the aircraft measurements during MILAGRO/INTEX-B are shown separately for DC-8 and C-130 in Figure

4. Data points below the 800 hPa pressure level were from observations at lower altitudes further away from the high-elevation MCMA. Values shown in Figure 4 are binned every 50 hPa vertically. The NO_x profiles are shown here in addition to O_3 and CO for the purpose of indicating the altitude range for the MCMA pollution outflow. The outflow occurs mainly at 600-800 hPa, corresponding to the high elevation of the Mexico City Basin (~750 hPa), as indicated by the enhanced O_3 , CO, and NO_x levels.

There are generally higher concentrations for all three pollutants, O_3 , CO, and NO_x at 500-800 hPa in the C-130 observations than those from the DC-8, while the later shows higher concentrations at 700-900 hPa. The higher concentrations in the C-130 observations at 500-800 hPa are due presumably to the proximity of the C-130 flights to the urban area of the MCMA (Figure 1) thus exposure to more pollution events. The higher concentrations in the DC-8 observations at 700-900 hPa likely indicate the export of MCMA pollution by the prevailing southwesterlies in the middle troposphere to downwind regions (e.g., near the Gulf of Mexico) where there is more DC-8 coverage, followed by subsidence (*Fast et al.* 2007).

As stated in the Section 2.2, TES averaging kernels were applied to *in situ* profiles for direct comparisons in previous TES validation studies with relaxed coincidence criteria. However, such coincidence criteria are generally inappropriate for a relatively small regional domain such as the one used in the present analysis (rectangle, Figure 1). During MILAGRO/INTEX-B there is only one coincident spiral profiling by the DC-8 for TES validation on March 12th (23°N and 98.5°W, ~450 km away from the MCMA). Validation results using profiles from that spiral have been reported by *Richard et al* (2008) and *Luo et al* (2007) and was cited in Section 2.2. In general, direct comparisons of absolute mixing ratios between TES and *in situ* data over the MCMA and its surrounding regions can be challenging primarily because of the scarcity of coincident TES

and aircraft profiles. We instead intend to investigate the spatial and temporal variabilities of O₃ and CO and the O₃-CO correlation as seen from TES and *in situ* data and GEOS-Chem results.

209

210

211

212

213

214

215

216

217

218

219

220

221

222

223

224

225

226

227

228

207

208

3.2. Vertical distributions of TES tropospheric O₃ and CO over the MCMA

Figure 5 shows individual TES vertical O₃ profile during MILAGRO/INTEX-B (gray dots) within the domain defined in Figure 1. The median TES vertical O₃ profile is also shown (data binned every 50 hPa). We focus our analysis on the 600-800 hPa pressure levels where most of the MCMA outflow occurs (Figure 4). Large enhancements and variations in O₃ are seen in several profiles. Such enhancements are not seen in the TES a priori (not shown). In addition, the mean DOFs is greater than 0.4 at 600 – 900 hPa, indicating that the TES O₃ profiles are not entirely determined by the a priori. Figure 6 shows the corresponding individual and monthly median vertical profiles of TES CO. Several profiles with enhanced CO concentrations at 500 – 800 hPa (> 130 ppbv) are not seen in the *a priori* (not shown). Also shown in Figures 5 and 6 are TES O₃ and CO profiles over unpolluted regions over the tropical Pacific adjacent to the west coast of Mexico (103°W ~ 107°W, 15°N – 25°N, black dots in Figures 5 and 6). The notably enhanced concentrations (> 70 ppbv for O₃ and >130 ppbv for CO, indicated by the gray dots) are only seen in the profiles over the MCMA and its downwind regions. This indicates that TES captures on a regional scale the enhancements in O₃ and CO as a result of the MCMA pollution outflow. Figure 7 shows the zonal mean of TES O₃ and CO in March 2006 over the domain defined in Figure 1 (rectangle). TES does not clearly show the pollution over the Mexico City Basin (~19N°) due mainly to the retrieval issues (see Section 2.2). The enhanced TES O₃ and CO

between 20 - 23N° is likely partially due to the MCMA pollution outflow transported by the southwesterly. This pattern is seen in TES data from other years (not shown).

231

232

233

234

235

236

237

238

239

240

241

242

243

244

245

246

247

248

249

250

251

229

230

3.3. O₃ and CO variabilities over the MCMA

Our analysis in this Section focuses on the variabilities of lower tropospheric O₃ and CO and to what degree they are captured by TES observations. More specifically, we examine the dayto-day and spatial variabilities of TES O₃ and CO in comparison with the *in situ* observations. Figure 8 shows time series of O₃ and CO from the *in situ* aircraft observations and TES retrievals during MILAGRO/INTEX-B. Values shown are for 600-800 hPa. The in situ data are shown without applying TES averaging kernels for reasons discussed in Section 3.1. GEOS-chem model results are also shown without TES averaging kernels applied. On average the application of TES averaging kernels to GEOS-Chem concentration fields makes little difference: < 1 ppbv for O₃ and < 6 ppbv for CO, respectively. We emphasize here that Figure 8 is intended for examining spatial and temporal varibilities of O₃ and CO instead of their absolute magnitudes among the three data sets. The *in situ* data in Figure 8 indicate nine strong pollution days: March 4th, 8th, 10th, 11th, 12th, 16th, 19th, 22nd, and 29th with O₃ and CO concentrations reaching 100 ppbv and 300 ppbv, respectively. Six of the pollution events were observed by C-130 while four were observed by DC-8, again reflecting presumably C-130's tighter geographical coverage over the MCMA. On four of the strong pollution days, March 4th, 11th, 22nd, and 29th, there were no concurrent TES observations or no TES coverage. Note that the ranges of O₃ and CO concentrations in Figure 8 on any individual day are indicative of the spatial variabilities of O₃ and CO considering the relatively large spatial coverage of the satellite and aircraft flight tracks (see Figure 1). Figure 8

shows that TES O₃ shows to a large extent both the spatial and day-to-day variabilities over the MCMA regions. For example, enhanced O₃ levels are seen in TES data on March 10th and 12th as in the *in situ* observations, as well as on March 25-27th when no in situ observations are available. TES CO, however, shows much less spatial and day-to-day variabilities compared with the *in situ* data.

GEOS-Chem results show little temporal variations in both O₃ and CO and significant negative biases: -15 ppbv for O₃ and -55 ppbv for CO in comparison with the *in situ* data. This may be in part due to the low CO emissions in the BRAVO inventory as NOx emissions in the same inventory was found to be underestimated (*Boersma et al.*, 2008). In addition, the model resolution at 2°x2.5° may be too coarse to capture the spatial variabilities seen in the *in situ* data. Further model analysis with finer horizontal resolutions would be helpful to understand the pollution and its outflow over the MCMA.

We take a closer look at the temporal mismatch between TES O₃ and CO data and the *in situ* observations. An example is shown in Figure 9 for March 10-12th, when both TES retrievals and the *in situ* observations were closest to the MCMA. There are large temporal and spatial variabilities in both O₃ and CO during the course of the aircraft flights. The peaks in the *in situ* observations occurred after 13:00 local time due to the development of strong nocturnal vertical inversions over the MCMA (*Velasco et al.*, 2007). TES data provide a snap shot at around 01:45 and 13:45 local time only, thus missing the intra-day temporal variability. However, TES data, especially O₃, do show substantial spatial variability during the short period of overpass (~20 minutes) over the domain defined in Figure 1, comparable to those in the corresponding DC-observations. TES CO shows consistently lower variability. The implication of low CO variability for calculated TES O₃-CO correlation will be discussed in the next section.

275

276

277

278

279

280

281

282

283

284

285

286

287

288

289

290

291

292

293

294

295

296

297

3.4. O₃-CO over the MCMA

As discussed in the previous sections, TES captured little of the CO day-to-day variability at 600-800 hPa during MILAGRO/INTEX-B (Figures 8 & 9). This would conceivably contribute to higher $\Delta O_3/\Delta CO$ enhancement ratios. We examine in this section the O_3 -CO correlations and slopes for 600-800 hPa derived from the aircraft measurements, TES retrievals, and GEOS-Chem results during MILAGRO/INTEX-B. The results are summarized in Table 1 and shown in Figure 10. All the data are averages on $2^{\circ} \times 2.5^{\circ}$ grids over the domain defined in Figure 1. The in situ aircraft observations during MILAGRO/INTEX-B show a strong O₃-CO correlation (correlation coefficient R = 0.78) and a positive $\Delta O_3/\Delta CO$ enhancement ratio of 0.28 mol mol⁻¹ for the domain defined in Figure 1. The regression results were derived using the reduced major axis method (Hirsh and Gilroy, 1984). The slope is within the range of those derived from summertime in situ measurements over the Eastern U.S. at the surface and in the lower troposphere: $\Delta O_3/\Delta CO = 0.2$ -0.4 mol mol⁻¹ with R = 0.7-0.9 (e.g., *Chin et al.*, 1994; Parrish et al., 1993). In comparison, slopes derived from the International Consortium on Atmospheric Transport and Transformation (ICARTT) aircraft campaigns in July-August 2004 (Fehsenfeld et al., 2006; Singh et al., 2006) are $\Delta O_3/\Delta CO = 0.31$ -0.44 mol mol⁻¹ with R = 0.5-0.67 in the lower troposphere over the Eastern U.S. (30°N – 50°N, 50°W – 100°W, and surface-600 hPa). Previously, Zhang et al. (2006) reported a slope of $\Delta O_3/\Delta CO = 0.72$ mol mol⁻¹ derived from in situ aircraft measurements during the same ICARTT campaign over the Eastern U.S. The difference is that Zhang et al. (2006) excluded fresh pollution plumes (NOx/NOy > 0.4 mol mol⁻¹, altitude < 3 km, and HCN > 500 pptv or CH₃CN > 225 pptv) and stratospheric influence (O₃/CO $> 1.25 \text{ mol mol}^{-1}$) within a smaller domain (30°N – 40°N, 70°W – 80°W, and 600 – 650 hPa). In

comparison, following *Zhang et al.* (2006) except without excluding fresh pollution and stratospheric influence, the $\Delta O_3/\Delta CO$ for the ICARTT campaigns is 0.55 mol mol⁻¹, significantly lower than that reported by *Zhang et al.* (2006).

TES O₃-CO during MILAGRO/INTEX-B shows a correlation coefficient of R = 0.5 and a slope of Δ O₃/ Δ CO = 0.43 mol mol⁻¹ for the domain defined in Figure 1. The slope is significantly larger than that derived from the *in situ* data (0.28 mol mol⁻¹). As a comparison, *Zhang et al.* (2006) derived a TES Δ O₃/ Δ CO enhancement ratio of 0.81 mol mol⁻¹ over the Eastern U.S. (30°–40°N, 70°–80°W) for July 2005. Their calculations were done for TES data at 618 hPa and within 10°×10° domains. *Zhang et al.* (2006) noted that their slope of 0.81 mol mol⁻¹ is comparable with the value of 0.72 mol mol⁻¹ derived from the ICARTT *in situ* data when excluding fresh plumes and stratospheric influence. It is however significantly higher than the value of 0.55 mol mol⁻¹ when not filtering out fresh plumes and stratospheric influence.

Whether fresh plumes and stratospheric influence should be filtered out in this type of comparisons between TES (and other space-borne instrument) versus aircraft observations remains a question. It is also important to note that a comparison between $\Delta O_3/\Delta CO$ slopes derived from TES retrievals and *in situ* aircraft observations may not be straightforward when the latter is not convoluted with TES averaging kernels. Model results, however, can be directly and straightforwardly compared with both *in situ* observations and TES data and therefore can be used as a transfer function between TES and *in situ* data. The GEOS-Chem $\Delta O_3/\Delta CO$ enhancement ratios during MILAGRO/INTEX-B with and without TES averaging kernels applied show similar results, 0.25 versus 0.30 mol mol⁻¹, respectively. The slopes are comparable with that derived from the *in situ* observations (0.28 mol mol⁻¹) but much lower that that derived

from TES data (0.43 mol mol⁻¹). The higher slopes derived from TES data could be partly attributed to the lack of variability and a low bias in TES CO in the lower troposphere as we discussed in previous sections.

4. Conclusions

320

321

322

323

324

325

326

327

328

329

330

331

332

333

334

335

336

337

338

339

We examined O₃, CO, and their relationships from TES tropospheric retrievals (v003), aircrafts observations, and GEOS-Chem model results over the Mexico City Metropolitan Area (MCMA) and its surrounding regions (15N°-30°N and 95°W-103°W) during the MILAGRO/INTEX-B campaigns. The MCMA pollution outflow occurs predominantly at 600-800 hPa as evident in enhanced concentrations in O₃ and CO in the aircraft observations. TES averaging kernels of O₃ and CO have generally higher sensitivity at 600 – 800 hPa, but there is poor sensitivity over the Mexico City Basin (~750 hPa) due to retrieval issues. Thus TES data are mostly sensitive to the MCMA pollution outflow rather than pollution over the MCMA itself. Direct comparison of absolute concentrations between *in situ* and TES data is difficult due to the spatiotemporal mismatches. Instead, we examined the spatial and temporal variability of TES O₃ and CO data and O₃-CO correlation. TES O₃ exhibits to a large degree the spatial and day-today variability as see in the *in situ* observations. There is significantly smaller spatiotemporal variability in addition to a low bias in TES lower tropospheric CO compared with in situ observations over the MCMA and its surrounding regions. Both likely contribute to the apparent overestimate of TES ΔO₃/ΔCO enhancement ratios during MILAGRO/INTEX-B (0.43 mol mol 1) compared with that from the *in situ* data (0.28 mol mol⁻¹). Thus the use of TES tropospheric O₃ and CO retrievals for mapping continental pollution outflow needs further examination.

341

342

340

Acknowledgements

This work is performed at the Jet Propulsion Laboratory (JPL), California Institute of Technology, under contract with the NASA. C. Shim and Q. Li were jointly supported by the JPL Research and Technology Development (R&TD) program, Human Resources Development Fund (HRDF), NASA Atmospheric Composition Modeling and Analysis program (ACMAP), and the TES project at JPL. The GEOS-Chem model is managed at Harvard University with support from the NASA Atmospheric Chemistry Modeling and Analysis Program. We thank Melody Avery for the *in situ* O₃ measurements. We thank Lin Zhang for very helpful discussions.

References

- Avery, M. A., Westberg, D. J., Fuelberg, H. E., *et al.*, Chemical transport across the ITCZ in the central Pacific during an El Ni no-Southern Oscillation cold phase event in March–April 1999, J. Geophys. Res., 106 (D23), 32 539–32 554, doi:10.1029/2001JD000728, 2001.
- Beer, R., TES on the Aura Mission: Scientific objectives, measurements and analysis overview, IEEE Trans. Geosci. Remote. Sens., 44, 1102 1105, 2006.
- Beer, R., Glbavich, T. A., and Rider, D. M., Tropospheric emission spectirometer for the Earth Observing System's Aura satellite, Appl, Opt, 40, 2356 2367, 2001.
- Benkovitz, C. M., Scholtz, M. T., Pacyna, J., *et al.*, Global gridded inventories of anthropogenic emissions for sulfur and nitrogen, J. Geophys. Res., 101 (D22), 29,239 29,253, 1996.
- Bey, I., Jacob, D. J., Yantosca, R. M., *et al.*, Global modeling of tropospheric chemistry with assimilated meteorology: Model description and evaluation, J. Geophys. Res., 106(D19), 23,073 23,096, 2001.
- Boersma, K. F., D. J. Jacob, E. J. Bucsela, *et al.*, Validation of OMI tropospheric NO₂ observations during INTEX-B and application to constrain NO_x emissions over the eastern United States and Mexico, Atmos. Environ., in press, 2008.
- Bowman, K. W., J. Worden, T. Steck, et al., Capturing time and vertical variability of tropospheric ozone: A study using TES nadir retrievals, J. Geophys. Res., 107(D23), 4723, doi:10.1029/2002JD002150, 2002.
- Bowman, K. W., Rodgers, C. D., Kulawik, S. S., *et al.*, Tropospheric emission spectrometer: Retrieval method and error analysis, IEEE Trans. Geosci. Remote. Sens., 44, 1297 1307, 2006.

- Brasseur, G. P., Hauglustaine, D. A., Walters, S, *et al.*, MOZART: A global chemical transport model for ozone and related chemical tracers, Part I: Model description, J. Geophys. Res., 103, 28,256 28,289, 1998.
- CAM, 1998 Preliminary Emissions Inventory; INE, 2000b, Almanaque de Datos y Tendencias de la Calidad del Aire en Ciudades Mexicanas, 2001.
- Clerbaux, C. Edwards, D. P., Deeter, M. *et al.*, Carbon monoxide pollution from cities and urban areas observed by the Terra/MOPITT mission, Geophys. Res. Lett., 35, L03817, doi:10.1029/2007GL032300.
- Chameides, W. L., David, D. D., Rodgers, M. O., *et al.*, Net ozone photochemical production over the eastern and central north Pacific as inferred from CTE/CITE 1 observations during fall 1983, J. Geosphys. Res., 92, 2131 2152, 1987.
- Chin, M., Jacob, D. J., Munger, J. W., *et al.* Relationship of ozone and carbon monoxide over North America, J. Geophys. Res., 99, 14,565 14,573, 1994.
- Duncan, B. N., Martin, R. V., Staudt A. C., *et al.*, Interannual and seasonal variability of biomass burning emissions constrained by satellite observations, J. Geophys. Res., 108(D2), 4100, doi:10.1029/2002JD002378, 2003.
- EPA, U. S., EPA Clearinghouse for inventories and emissions factors: 1999

 National Emission Inventory Documentation and Data Final Version 3.0,

 http://www.epa.gov/ttn/chief/net/1999inventory.html, 2004.
- Fast, J. D., B. de. Foy, F. Acevedo Rosas, *et al.*, A meteorological overview of the MILAGRO field campaigns, Atmos. Chem. Phys., 7, 2233-2257, 2007.

- Fehsenfeld, F. C. et al., International Consortium for Atmospheric Research on Transport and Transformation (ICARTT): North America to Europe Overview of the 2004 summer field study, J. Geosphys. Res., 111, D23S01, doi:10.1029/2006JD007829 (2006).
- Fishman, J., and Seiler, W., Correlative nature of ozone and carbon monoxide in the troposphere: Implications for the tropospheric ozone budget, J. Geosphys. Res., 88, 3662 3670, 1983.
- Fuchs, R. J., Brennan, E., Chamie, J., *et al.*, Mega-city growth and the future, United Nations university press, Tokyo, 1994.
- Guenther, A.B., Karl, T., Harley, P., *et al.*, Estimates of global terrestrial isoprene emissions using MEGAN (Model of Emissions of Gases and Aerosols from Nature), Atmos. Chem. Phys, 6, 3181-3210, 2006.
- Hirsch, R. M., and Gilroy, E. J., Methods of fitting a straight line to data: Examples in water resources, Water Resour. Bull., 20,705 711, 1984.
- Hudman, R. C., Jacob, D. J., Turquety, S., *et al*, Surface and lightning sources of nitrogen oxides over the United States: Magnitudes, chemical evolution, and outflow, J. Geophys. Res., 112, D12S05, doi:10.1029/2006JD007912, 2007.
- Jacob, D. J., Crawford, J. H., Kleb, M. M., *et al.*, Transport and Chemical Evolution over the Pacific (TRACE-P) aircraft mission: Design, execution, and first results, J. Geophys. Res., 108(D20), 9000, doi:10.1029/2002JD003276, 2003.
- Jones, D. B. A., Bowman, K. W., Palmer, P. I., et al., Potential of observations from the Tropospheric Emission Spectrometer to continental sources of carbon monoxide, J. Geophys. Res., 108(D24), 4789, doi: 10.1029/2003JD003702, 2003.
- Jourdain, L. and Hauglustaine, D. A., The global distribution of lightning NOx simulated on-line in a general circulation model, Phys. Chem. Earth, C26, 585–591, 2001.

- Jourdain, L., Worden, H. M., Worden, J. R., *et al.*, Tropospheric vertical distribution of tropical Atlantic ozone observed by TES during the northern African biomass burning season, Geophys. Res. Lett., 34, L04810, doi:10.1029/2006GL028284, 2007.
- Kuhns, H., Knipping, E. M., Vukovich J. M., *et al.*, Development of a United States-Mexico Emissions Inventory for the Big Bend Regional Aerosol and Visibility Observational (BRAVO) Study, J. Air. Waste Manag. Assoc., 55(5): 677-92, 2005.
- Logan, J. A., Prather, M. J., Wofsy, S. C., *et al.*, Tropospheric Chemistry: A global perspective, J. Geophys. Res., 86, 7210-7254, 1981.
- Luo M., Rinsland, C. P., Rodgers, C. D., *et al.*, Comparison of carbon monoxide measurements by TES and MOPITT: the influence of a priori data and instrument characteristics on nadir atmospheric species retrievals, J. Geophys. Res., 112, D09303, doi:10.1029/2006JD007663, 2007a.
- Luo M., Rinsland, C., Fisher, B., *et al.*, TES carbon monoxide validation with DACOM aircraft measurements during INTEX-B 2006, J. Geophys. Res., in press, 2007b.
- Madronich S., Flocke, F. Orlando, J., *et al.*, MIRAGE-Mex: Mexico City Pollution Outflow Experiment: Science Overview Document, http://mirage-mex.acd.ucar.edu/Documents/MIRAGE-Mex SOD 040324.pdf, 2004.
- Mage, D., Ozolins, G., Peterson, P., *et al.*, Urban air pollution in megacities of the world, Atmos. Environ, 30(5), 681-686, 1996.
- Massie S. T., Gille J. C., Edwards, D. P., *et al.*, Satellite Observations of aerosol and CO over Mexico city, Atmos. Environ., 40, 6019 6031, 2006.
- Molina, L.T., and Molina, M. J., Air Quality in the Mexico Megacity, Kluwer Academic Publishers, Norwell, M.A., 2002.

- Molina, M. J. and Molina, M. J., Critical Review: Megacities and Atmospheric Pollution. J. Air & Waste Manage. Assoc., 54, 6, 644 680, 2004.
- .Olivier, J.G.J. and Berdowski, J. J. M., Global emissions sources and sinks. In: Berdowski, J., Guicherit, R. and B.J. Heij (eds.) "The Climate System", pp. 33-78. A.A. Balkema Publishers/Swets & Zeitlinger Publishers, Lisse, The Netherlands. ISBN 90 5809 255 0, 2001.
- Osterman, G., Bowman, K., Cady-Pereira, K., *et al.*, Tropospheric emission spectrometer (TES) validation report, JPL D33192, version 2.0 (available at http://tes.jpl.nasa.gov/docsLinks/documents.ctm), 2006.
- Park, R. J., Jacob, D. J., Field, B. D., *et al.*, Natural and transboundary pollution influences on sulfate-nitrate-ammonium aerosols in the United States: Implications for policy, J. Geophys. Res., 109, D15204, doi:10.1029/2003JD004473, 2004.
- Parrish, D. D., Holloway, J. S., Trainer, M., *et al.*, Export of North American ozone pollution to the North Atlantic Ocean, Science, 259, 1436 1439, 1993.
- Richards, N. A. D., Q. Li, K. W. Bowman, *et al.*, Assimilation of TES CO into a global CTM: first results, Atmos. Chem. Phys. Discuss., 6, 11727-11743, 2006.
- Richards, N. A. D., Osterman, G. B., Browell, E. V., *et al.*, Validation of Tropospheric Emission Spectrometer (TES) Ozone Profiles with Aircraft Observations During INTEX-B, J. Geophys. Res., 113, D16S29, doi:10.1029/2007JD008815, 2008.
- Rinsland, C. P., M Luo, M. W. Shephard, et al, Tropospheric Emission Spectrometer (TES) and Atmospheric Chemistry Experiment (ACE) measurements of tropospheric chemistry in tropical southeast Asia during a moderate El Niño in 2006, J Quant Spectrosc Radiat Transfer, December 2007, in press.

- Rogers, C. D., Inverse Methods for Atmospheric Sounding: Theory and Practice, World Sci., River Edge, N. J., 2000.
- Rogers, C. D., and B. J. Conner, Intercomparison of remote sounding instruments, J. Geophys. Res., 108(D3), 4116, doi:10.1029/2002JD002299.
- Sachse, G. W., G. F. Hill, L. O. Wade et al., Fast-response, high-precision carbon monoxide sensor using a tunable diode laser absorption technique, J. Geophys. Res. 92, 2071 2081, 1987.
- Singh, H. B., Brune, W. H., Crawford, J. H. et al., Overview of the summer 2004

 Intercontinental Chemical Transport Experiment North America (INTEX-A), J.

 Geophys. Res., 111, D24S01, doi:10.1029/2006JD007905 (2006).
- Singh, H. B., Brune, W. H., Crawford, J. H., *et al.*, An overview of the INTEX-B campaign: Transport and transformation of pollutants over the Pacific and the Gulf of Mexico, Atmos. Chem. Phys. Discuss, in preparation, 2008.
- Velasco, E., C. Marquez, E. Bueno, *et al*, Vertical distribution of ozone and VOCs in the low boundary layer of Mexico City, Atmospheric Chemistry and Physics Discussions, 7, 12751 12779, 2007.
- Worden, H. M., Logan, J. A., Worden, J. R., *et al.*, Comparisons of Tropospheric Emission Spectrometer (TES) ozone profiles to ozonesodes: methods and initial results, J. Geophys. Res., 112, D03309, doi:10.1029/2006JD007258, 2007.
- Worden, J., Kulawik, S. S., Shephard, M. W., *et al.*, Predicted errors of tropospheric emission spectrometer nadir retrievals from spectral window selection, J. Geophys. Res., 109, No. D9, D09308, 10.1029/2004JD004522, 2004.

Zhang, L., Jacob, D. J., Bowman, K. W., *et al.*, Ozone-CO correlations determined by the TES satellite instrument in continental outflow regions, Geophys. Res. Lett., 33, L18804, doi:10.1029/2006GL026399, 2006.

Table

Table 1. O₃ – CO correlation coefficient and slope (Δ O₃/ Δ CO)

Location		Time period	Pressure	r	$\Delta O_3/\Delta CO$	
MCMA	in situ	March 2006	600 – 800 hPa	0.78 (0.69) ^a	0.28±0.07	0.36±0.05 ^b
	TES			0.5	0.43 ± 0.1	
	GC/AK ^c			0.58	0.25 ± 0.06	
	GC-raw ^d			0.26	0.3±0.15	
Eastern U.S. (TES ^e)		July 2005	618 hPa	0.53	0.81	Zhang <i>et al.</i> , (2006)
Eastern U.S. ^f		July- September, 2004	600 hPa – surface	0.5 - 0.7	0.31 - 0.44	ICARTT
Western Pacific		Feburary- March, 2001	600 hPa – surface	0.6	0.15	TRACE-P
Eastern U.S.		June – August, 1988 - 1991	surface	0.7 - 0.9	0.2 - 0.4	Chin <i>et al</i> . (1994)
Sable Island		July – September, 1991	surface	0.82	~0.3	Parrish <i>et al.</i> (1993)

All MCMA data were averaged onto 2°×2.5° grids over the MCMA (shown in Figure 1).

^aThe correlation coefficient inside parenthesis indicates the result excluding the fresh pollution plumes and stratospheric O₃ influence according to Hudman *et al.*, (2007)

^bThe $\Delta O_3/\Delta CO$ derived from the same data as (a).

^cGEOS-Chem results with TES averaging kernels applied.

dGEOS-Chem results without applying TES averaging kernels.

^eTES version 1(V001) data

^fThe data were selected from DC-8 between $30N^{\circ} - 50N^{\circ}$ and $50W^{\circ} - 100W^{\circ}$ without the filtering as (a).

Figures

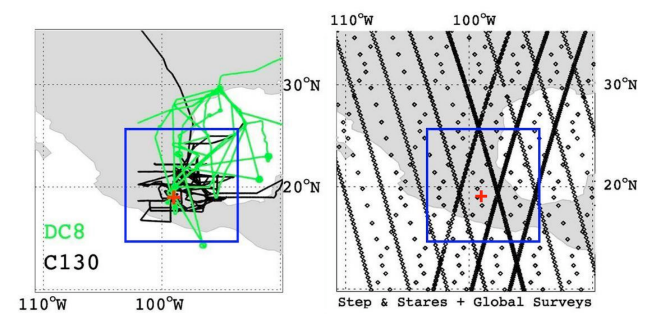


Figure 1. Flight tracks of the C-130 (black) and DC-8 (green) aircrafts during the MILAGRO/INTEX-B (phase I) campaigns in March 2006 (left panel). Also shown are TES orbital tracks (eight Step & Stares and four Global Surveys) during the same time period (right panel). The blue rectangle indicates the domain for analysis presented in this study.

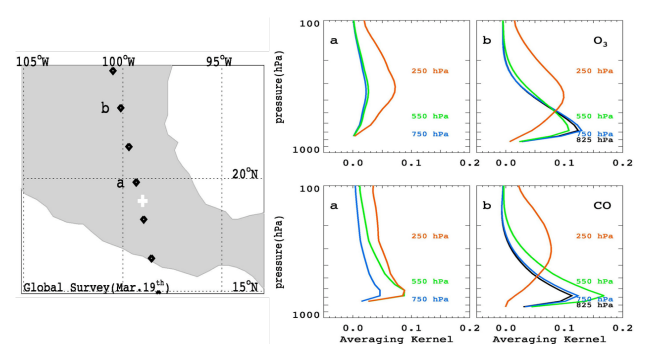


Figure 2. TES averaging kernels of O₃ (upper right panels) and CO (lower right panels) for Global Surveys closest to MCMA (white cross on the left panel) during MILAGRO/INTEX-B campaign (March 19th, 2006). The location "a" (left panel) denotes the closest TES retrieval to MCMA (~95 km from MCMA) and "b" (left panel) denotes TES retrieval where its distance is ~470 km from MCMA. Averaging kernels for different pressure levels are color-coded.

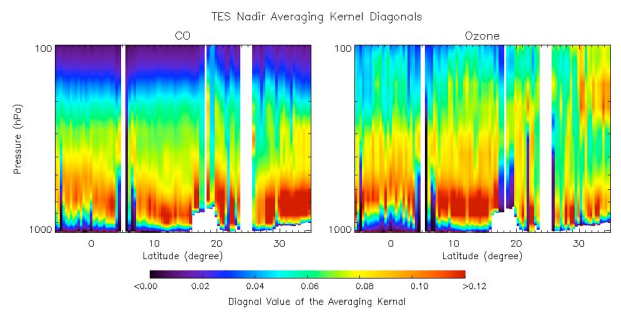


Figure 3. The TES Nadir Averaging Kernel Diagonals for CO (left) and O₃ (right) along the special observations in Figure 2.

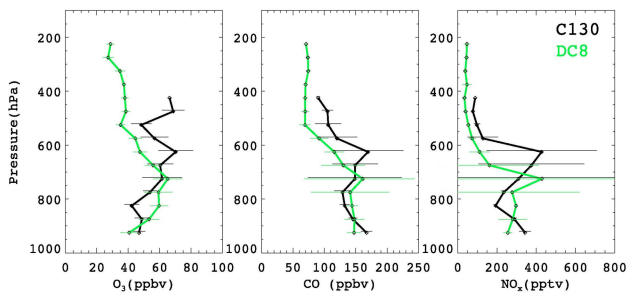


Figure 4. The median vertical profiles of O₃ (left panel), CO (middle panel), and NOx (right panel) binned 50 hPa from C130 (black) and DC8 (green) during MILAGRO/INTEX-B (Phase I). Standard deviations are shown for each pressure level.

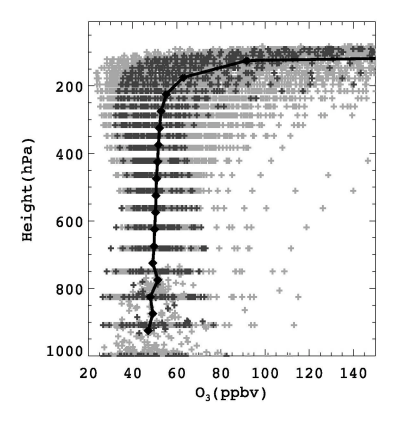


Figure 5. The collected TES vertical profiles of O_3 (gray dots) over MCMA and surrounding regions (15–25°N and 95 - 103°W) in March 2006. The black dots indicate the vertical profiles over the unpolluted region (15–25N° and 103 ~ 107W°). The black solid line indicates the median profiles binned 50 hPa.

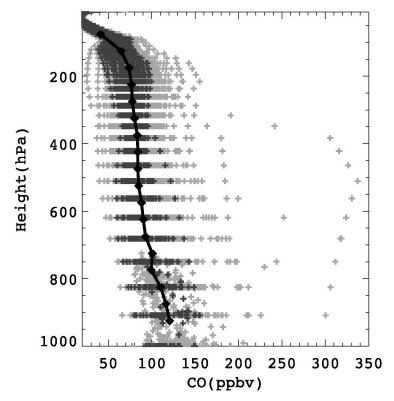


Figure 6. Same as Figure 5, but for CO.

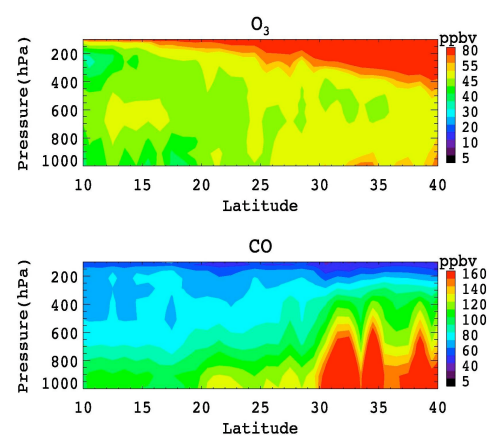


Figure 7. Zonal mean concentration of TES O_3 (upper) and CO (lower) between $(95W^\circ - 103W^\circ)$ in March 2006.

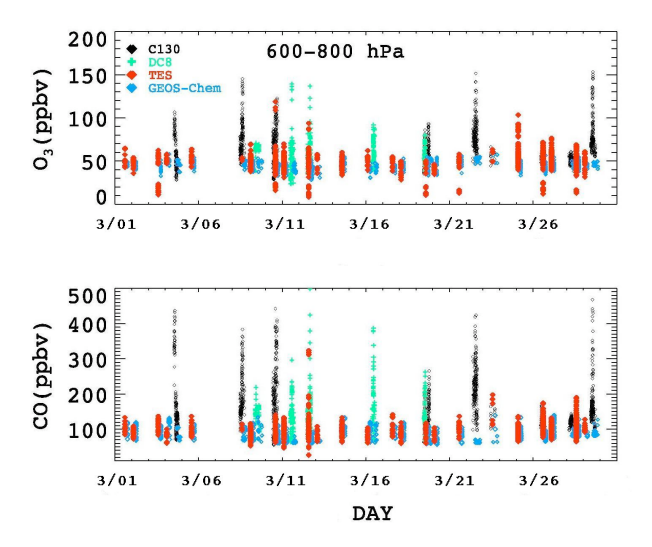


Figure 8. Time series of individual O_3 (upper panel) and CO (lower panel) data at 600 - 800 hPa over MCMA in March 2006. *in-situ* observations from C130 and DC8, TES retrievals, and GEOS-Chem results are indicated by black diamonds and green cross, red, and blue diamonds, respectively. GEOS-Chem results are sampled along TES orbital and/or aircraft flight tracks.

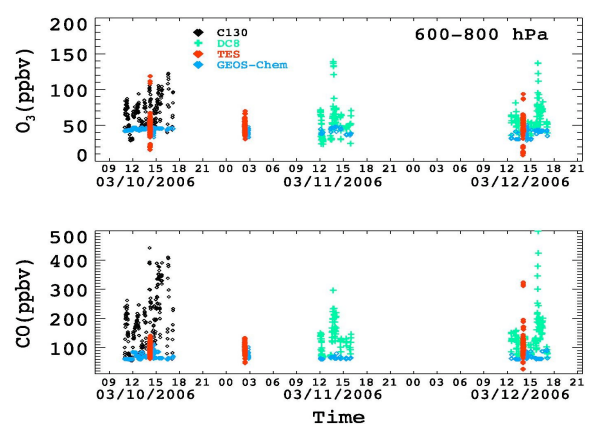


Figure 9. Same as Figure 7, but with diurnal variations for March 10^{th} - 12^{th} when both the *in-situ* and TES samplings (Step & Stares) are closest to the MCMA (within $\sim 200 \text{ km}$).

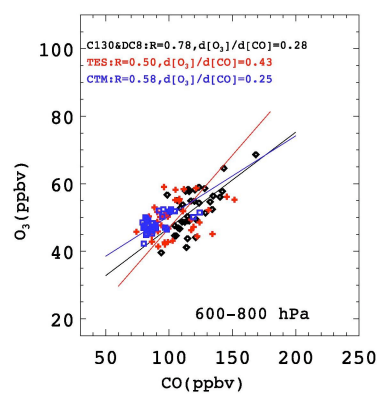


Figure 10. CO-O₃ relationships at 600 - 800 hPa during MILAGRO/INTEX-B (Phase I). *in-situ* and TES data were averaged onto $2^{\circ} \times 2.5^{\circ}$ grids over the MCMA and surrounding region. Black diamonds - aircraft measurements; red crosses - TES; blue rectangles - GEOS-Chem results sampled along TES orbital tracks with TES averaging kernels applied.